principle.

1 TRANSITION METAL COMPOUND

The present invention relates to a zwitterionic, neutral transition metal compound which can be advantageously used for the polymerization of olefins. In this way, the use 5 of aluminoxanes such as methylaluminoxane (MAO) as cocatalyst can be omitted while still achieving high catalyst activities.

The role of cationic 14-electron complexes of the formula R₂M⁺ (M=Ti, Zr, Hf) in Ziegler-Natta polymerization 10 using metallocenes is generally recognized (M. Bochmann, Nachr. Chem. Lab. Techn. 1993, 41, 1220).

While MAO, as the hitherto most effective cocatalyst. has to be used in a high excess, the synthesis of cationic alkyl complexes of the formula R₃M+ (M=Ti, Zr, Hf) 15 provides a route to MAO-free catalysts, some of which have comparable activity.

Cationic alkyl complexes can be prepared by

a) oxidation of metallocene-dialkyl complexes with, for example, AgBPh₄ or [Cp₂Fe][BPh₄],

b) by protolysis of metallocene-alkyl compounds with, for example, weakly acid ammonium salts of the very stable, nonbasic tetra(pentafluorophenyl)borate anion (e.g. $[PhMe_2NH]^+[B(C_6F_5)_4]^-)$ or by

c) abstraction of an alkyl group from metallocene-alkyl 25 compounds by means of strong Lewis acids. Lewis acids which can be used here are salts (Ph₂C+BR₄-) or strong neutral Lewis acids such as $B(C_6F_5)_3$.

J. Organomet. Chem. 1970, 22, 659, describes a reaction of tetramethyltitanium with triphenylborane or tribenzylbo- 30

J. Am. Chem. Soc. 1991, 113, 3623, describes the synthesis of "cation-like" metallocene polymerization catalysts which are prepared by alkyl abstraction from metallocenealkyl compounds using tris(pentafluorophenyl)borane. The 35 crystal structure of [1,2-(CH₃)₂C₅H₃]₂-ZrCH3]⁺[CH₃B (C₆F₅)₃] shows a salt-like structure with weak coordination of the CH₃ group of the borate anion to the metal center. EP 427,697 claims this synthetic principle and a corresponding catalyst system comprising a neutral metallocene species 40 (eg. Cp2ZrMe2), a Lewis acid (eg. B(C6F5)3) and aluminum alkyls. EP 520,732 claims a process for preparing salts of the general formula LMX⁺XA⁻ according to the abovedescribed principle.

EP 558,158 claims zwitterionic catalyst systems prepared 45 from metallocene-dialkyl compounds and salts of the formula [R3NH]+[BPh4]. Reaction of such a salt with Cp*2ZrMe2 generates, by means of protolysis with elimination of methane, an intermediate zirconocene-methyl cation which reacts, after C-H activation of a tetraphenylbo- 50 rate carbon-hydrogen bond and renewed elimination of methane, to give the zwitterion Cp*2Zr+-(m-C6H4)-B-Ph3. The Zr atom is here covalently bonded to a carbon atom of the phenyl ring and is stabilized via an agostic hydrogen bond. According to this reaction principle, protolysis of a 55 metallocene-dialkyl species with a perfluorinated [R3NH]+ [B(C₆F₅)₄] salt in the first step likewise forms a cationic species, with the subsequent reaction (C-H activation) to give zwitterionic complexes (i.e. the metal atom is covalently bonded to the "former" anion) not being possible. 60 This process also uses metallocenes CP₂MR₂ in which the alkyl radicals R are cyclically bonded to one another, for example Cp₂Zr(2,3-dimethyl-1,3-butadiene). Salts of the formula [Cp₂Zr-R-RH]+[B(C₆F₅)₄] are formed after protonolysis.

U.S. Pat. No. 5,198,401 claims corresponding systems in which dimethylanilinium salts having perfluorinated tetraphenylborate anions are used. This reference also uses metallocenes Cp₂MR₂ in which the alkyl radicals R are cyclically bonded to one another, for example Cp₂Zr-(2,3dimethyl-1,3-butadiene). After protonolysis, salts of the formula $[Cp_2Zr-R-RH]^+[B(C_6F_5)_4]^-$ are likewise formed. EP 277,003, EP 277,004, EP 495,375 and WO 91/14713 claim systems according to a similar process

The processes described for preparing the cationic systems of the formula [R₃M]⁺[BR₄]⁻ (M=Ti, Zr, Hf) have the disadvantage that the cationizing reagents [R₂NH+BR₄] are sometimes complicated to synthesize and costly. In addition, there is the problem that an amine R₃N is formed from the ammonium salt after protonolysis, which amine, in the case of sterically unhindered metal centers, can coordinate to the strong Lewis acid R₃M⁺ cation (U.S. Pat. No. 5,198,401) and thus leads to low polymerization activities.

The zwitterionic complexes of the structure Cp2Zr+-m-C₆H₄B⁻Ph₃ have the disadvantage that the starting compounds are expensive and complicated to synthesize and have low polymerization activity.

Owing to their salt-like character, cationic systems of the structure [Cp₂MR]⁺[RB(C₆F₅)₃] have very high hydrolysis sensitivities and can be used on an industrial scale only to a limited extent. The activities observed for these systems are low owing to the abovementioned stability problems and the presumably sometimes strong coordination of the alkyl group of the borate anion to the metal center.

It is an object of the invention to find a transition metal compound which avoids the disadvantages of the prior art.

It has now been found that this object can be achieved by specific zwitterionic transition metal compounds.

The present invention accordingly provides a zwitterionic transition metal compound of the formula I

$$\begin{array}{c}
 & X \\
 & X \\
 & AR^{1} \\
 & AR^{1}
\end{array}$$
(I)

where

L are identical or different and are each a π-ligand or an electron donor, n is equal to 1, 2, 3 or 4,

M is a metal atom of group IIIb, IVb, Vb or VIb of the Periodic Table of the Elements,

X is a heteroatom or a hydrocarbon group having 1-40 carbon atoms,

B is a hydrocarbon group having 1-40 carbon atoms,

A is a metal atom of group Ib, IIb, IIIa, IIIb, IVa, Va, Vb, VIb, VIIb or VIIIb of the Periodic Table of the Elements,

R¹ are identical or different and are each a perhalogenated C₁-C₄₀-hydrocarbon radical, and m is equal to 1, 2, 3, 4 or 5.

The metal atoms M and A bonded to one another by covalent bonds via the structural elements X and B. If X is an allyl unit, the bonding of X to the metal atom X can be a σ -allyl or a π -allyl bond. π -Ligands are preferably unsubstituted or substituted cyclopentadienyl groups such as 2-methylcyclopentadienyl, indenyl, 2-methylindenyl, 2-methyl-4-phenylindenyl, 2-methyl-4,5-benzoindenyl, 2-methyl-4,6-diisopropylindenyl, 4,5-benzoindenyl, fluorenyl, 4,7-tert-butylfluorenyl or 2-methyl-4-(2-pyridyl) 65 indenyl.

For the purposes of the present invention, an electron donor is an atom of group IVa, Va, VIa or VIIa of the

Periodic Table of the Elements, which can bear substituents such as C_1 – C_{20} -hydrocarbon groups. Preference is given to O, NR_2^4 , NR_3^4 , NR_3^4 , PR_2^4 , PR_3^4 , PR_3^4 , PR_4^4 , PR_5^4 , $PR_5^$

Two radicals L can be linked to one another via a bridge 5

The bridge Z is preferably

=BR₂, -AIR², -Ge-, -O-, -S-, =SO, =SO₂, -NR₂, =CO, =PR² or =P(O)R², where R² and R³ are 25 identical or different and are each a hydrogen atom, a halogen atom, a C_1 - C_2 -alkyl group, a C_1 - C_1 -fluoroalkyl group, a C_6 - C_{10} -fluoroaryl group, a C_6 - C_{10} -alkeyl group, a C_6 - C_{10} -alkeyl group, a C_7 - C_{40} -arylalkyl group, a C_7 - C_{40} - arylalkyl group, a C_7 - C_{40} - $C_$ alkylaryl group, a C8-C40-arylalkenyl group and x is a number from zero to 18, or R2 and R3 together with the atoms-connecting them form one or more rings and R2 or/and R³ can be bonded to L and M² is silicon, germanium

Z can also link two or more identical or different groups L_mM⁺XBA⁻R¹_m with one another.

For the purposes of the present invention, a heteroatom is any atom of the Periodic Table of the Elements with the exceptions of carbon and hydrogen. Preference is given to 40 O, S and N.

Hydrocarbon groups X and B can be saturated or unsaturated, linear or branched, eg. a C1-C20-alkyl group, a C_6 - C_{14} -aryl group, a C_2 - C_{10} -alkenyl group, a C_7 - C_{40} -arylalkyl group, a C_7 - C_{40} -alkylaryl group or a C_8 - C_{40} arylalkenyl group. Preference is given to substituted or unsubstituted alkyl groups which can also have aromatic structural elements.

Preference is given to n=1 when M is an element of group IIIb of the Periodic Table of the Elements; n=2 when M is 50 a metal atom of group IVb of the Periodic Table of the Elements; n=3 when M is an element of group Vb of the Periodic Table of the Elements; and n=4 when M is an element of group VIb of the Periodic Table of the Elements.

R¹ is a C₁-C₄₀-hydrocarbon radical which is perhaloge- 55 dimethylsilanediylbis(2-methyl-4,6-diisopropylindenyl)Zr⁺ nated with halogen such as fluorine, chlorine, bromine or iodine, in particular a perhalogenated C1-C30-alkyl group such as trifluoromethyl, pentachloroethyl, heptafluoroisopropyl, or monofluoroisobutyl or a perhalogenated C₆-C₃₀-aryl group such as pentafluorophenyl, 60 heptachloronaphthyl, heptafluoronaphthyl or heptafluorotolyl.

Particular preference is given to compounds of the formula I, where

the Elements, such as titanium, zirconium or hafnium, n is equal to 2,

L are identical or different and are each a substituted or unsubstituted cyclopentadienyl group, where two radicals L can be linked to one another via a bridge Z and

Z is CR²R³ or SiR²R³ or a unit Si—(CR²R³)_x—Si which links two fragments $L_nM^+XBA-R^1_m$ with one another, where x is an integer from 0 to 10, preferably x=6,

X and B together form a three-membered to fivemembered (C3-C5)-alkyl chain which is saturated or unsaturated and can be substituted by C1-C20hydrocarbon radicals,

A is a metal of group Ib, IIb, IIIa, IVa, Va, Vb of the Periodic Table of the Elements,

R1 are identical or different and are each a perfluorinated alkyl or aryl group having from 1 to 20 carbon atoms

m is equal to 2, 3 or 4.

Very particular preference is given to compounds of the formula I, where

M is zirconium,

n is equal to 2,

L are identical or different and are each a substituted cyclopentadienyl group, where two radicals L are bonded to one another via a bridge Z, where Z is CR2R3 or SiR²R³,

X and B together form an unsaturated four-membered (C₄)-alkyl chain whose hydrogen atoms can also be replaced by C₁-C₂₀-alkyl groups,

A is a boron atom,

R1 are identical and are each a pentafluorophenyl group (C_6F_5) and

m is equal to 3.

Examples of compounds of the invention are: bis(cyclopentadienyl)Zr+CH₂CHCHCH₂B-(C₆F₅)₃; bis(methylcyclopentadienyl)Zr+C2CHCHCH2B-(C6F5)3; bis(n-butylcyclopentadienyl)Zr+CH₂CHCHCH₂B-(C₆F₅)₃; bisindenylZr⁺CH₂CHCHCH₂B⁻(C₆F₅)₃;

(tert-butylamido)dimethyl(tetramethyl-η⁵cyclopentadienyl)silaneZr+CH2CHCHCH2B-(C6F5)3; bis(2-methylbenzoindenyl) $Zr^+CH_2CHCHCH_2B^-(C_6F_5)_3$; dimethylsilanediylbis(2-methylindenyl) Zr^+ $CH_2CHCHCH_2B^-(C_6F_5)_3;$

dimethylsilanediylbisindenylZr+CH2CHCHCH2B-(C6F5)3; dimethylsilanediylbis(2-methylbenzoindenyl)Zr+ $CH_2CHCHCH_2B^-(C_6F_5)_3;$

dimethylsilanediyl(2-methylbenzoindenyl)(2-methylindenyl)Zr+CH₂CHCHCH₂B-(C₆F₅)₃;

dimethylsilanediyl(2-methylbenzoindenyl)(2-methyl-4phenylindenyl)Zr⁺CH₂ CHCHCH₂B (C₆F₅)₃;

dimethylsilanediyl(2-methylindenyl)(4-phenylindenyl)Zr+ $CH_2CHCHCH_2B^-(C_6F_5)_3$;

dimethylsilanediylbis(2-methyl-4-phenylindenyl)Zr+ $CH_2CHCHCH_2B^-(C_6F_5)_3;$

 $CH_2CHCHCH_2B^-(C_6F_5)_3;$

dimethylsilanediylbis(2-methyl-4-naphthylindenyl)Zr+ CH2CHCHCH2B-(C6F5)3;

isopropylidene(cyclopentadienyl)(fluorenyl)Zr+ $CH_2CHCHCH_2B^-(C_6F_5)_3;$

isopropylidene(cyclopentadienyl)(indenyl)Zr+ $CH_2CHCHCH_2B^-(C_6F_5)_3;$

[4- η^5 -cyclopentadienyl-4,7,7-trimethyl-(η^5 -4,5,6,7-tetrahydroindenyl)Zr+CH₂CHCHCH₂B-(C₆F₅)₃;

M is a metal atom of group IVb of the Periodic Table of 65 dimethylsilanediylbis(2-methylindenyl)Zr $OCH_2CH_2CH_2B^-(C_6F_5)_3;$

dimethylsilanediylbisindenylZr+OCH2CH2CH2B-(C6F5)3;

dimethylsilanediylbis(2-methylbenzoindenyl)Zr⁺ OCH₂CH₂B-(C₆F₅)₃;

dimethylsilanediyl(2-methylbenzoindenyl)(2methylindenyl)Zr+OCH₂CH₂CH₂B-(C₆F₅)₃;

dimethylsilanediyl(2-methylbenzoindenyl)(2-methyl-4- 5 phenylindenyl)Zr*OCH₂CH₂CH₂B⁻(C₆F₅)₃;

dimethylsilanediyl(2-methylindenyl)(4-phenylindenyl)Zr+ OCH₂CH₂B-(C₆F₅)₃;

dimethylsilanediylbis(2-methyl-4-phenylindenyl)Zr⁺ OCH₂CH₂B⁻(C₆F₅)₃;

dimethylsilanediylbis(2-methyl-4,6-diisopropylindenyl)Zr⁺ OCH₂CH₂CH₂B⁻(C₆F₅)₃;

dimethylsilanediylbis(2-methylindenyl)Zr+ CH₂CHCHCH₂B-(CF₃)₃;

dimethylsilanediylbisindenylZr+CH₂CHCHCH₂B-(CF₃)₃; dimethylsilanediylbis(2-methylbenzoindenyl)Zr+ CH₂CHCHCH₂B-(CF₃)₃;

dimethylsilanediyl(2-methylbenzoindenyl)(2-methylindenyl)Zr+CH₂CHCHCH₂B-(CF₃)₃;

dimethylsilanediyl(2-methylbenzoindenyl)(2-methyl-4-20 phenylindenyl)Zr+CH₂CHCHCH₂B-(CF₃)₃;

dimethylsilanediyl(2-methylindenyl)(4-phenylindenyl)Zr* CH2CHCH2B*(CF₃)₃;

dimethylsilanediylbis(2-methyl-4-phenylindenyl)Zr+ CH2CHCH2B-(CF3)3;

dimethylsilanediylbis(2-methyl-4,6-diisopropylindenyl)Zr* CH₂CHCHCH₂B⁻(CF₃)₃;

dimethylsilanediylbis(2-methyl-4-naphthylindenyl)Zr+ CH2CHCH2B-(CF3)3;

dimethylsilanediylbis(2-methylindenyl)Zr+CH₂C(CH₃)C 30 (CH₃)CH₂B-(CF₃)₃;

dimethylsilanediylbisindenylZr+CH₂C(CH₃)C(CH₃)CH₂B-(CF₃)₃;

dimethylsilanediylbis(2-methylbenzoindenyl)Zr+CH₂C (CH₃)C(CH₃)CH₂B-(CF₃)₃;

dimethylsilanediyl(2-methylbenzoindenyl)(2methylindenyl)Zr+CH₂C(CH₃)C(CH₃)CH₂B-(CF₃)₃;

dimethylsilanediyl(2-methylbenzoindenyl)(2-methyl-4phenylindenyl)Zr⁺CH₂C(CH₃)C(CH₃)CH₂B⁻(CF₃)₃;

dimethylsilanediyl(2-methylindenyl)(4-phenylindenyl)Zr⁺ 40 CH₂C(CH₃)C(CH₃)CH₂B⁻(CF₃)₃;

dimethylsilanediylbis(2-methyl-4-phenylindenyl)Zr+CH₂C (CH₃)C(CH₃)CH₂B-(CF₃)₃;

dimethylsilanediylbis(2-methyl-4,6-diisopropylindenyl)Zr+CH₂C(CH₃)C(CH₃)CH₂B⁻(CF₃)₃;

dimethylsilanediylbis(2-methyl-4-naphthylindenyl)Zr⁺ CH₂C(CH₃)C(CH₃)CH₂B⁻(CF₃)₃;

methylphenylmethylene(fluorenyl)(cyclopentadienyl)Zr+ CH,CHCHCH₂B-(C₆F₅)₃;

diphenylmethylene(fluorenyl)(cyclopentadienyl)Zr⁺ 50 CH₂CHCHCH₂B⁻(C₆F₅)₃;

isopropylidene(3-methylcyclopentadienyl)(fluorenyl)Zr+CH2CHCH2B-(C₆F₅)₃;

dimethylsilanediyl(3-tert-butylcyclopentadienyl)(fluorenyl)
Zr+CH₂CHCHCH₂B-(C₆F₅)₃;

diphenylsilanediyl(3-(trimethylsilyl)cyclopentadienyl) (fluorenyl)Zr+CH₂CHCHCH₂B-(C₆F₅)₃;

phenylmethylsilanediylbis(2-methylindenyl)Zr* CH₂CHCHCH₂B⁻(C₆F₅)₃;

phenylmethylsilanediylbisindenyl $Zr^+CH_2CHCHCH_2B^-$ 60 $(C_6F_5)_3$;

phenylmethylsilanediylbis(2-methyl-4,5-benzoindenyl)Zr⁺CH₂CHCHCH₂B⁻(C₆F₅)₃;

phenylmethylsilanediyl(2-methyl-4,5-benzoindenyl)(2methylindenyl)Zr⁺CH₂CHCHCH₂B⁻(C₆F₅)₃;

phenylmethylsilanediyl(2-methyl-4,5-benzoindenyl)(2-methyl-4-phenylindenyl)Zr+CH2CHCHCH2B+(C₆F₈)₃;

phenylmethylsilanediyl(2-methylindenyl)(4-phenylindenyl) $Zr^+CH_2CHCHCH_2B^-(C_6F_5)_3$;

phenylmethylsilanediylbis(2-methyl-4-phenylindenyl)Zr+ CH₂CHCHCH₂B⁻(C₆F₅)₃;

phenylmethylsilanediylbis(2-ethyl-4-phenylindenyl)Zr⁺ CH₂CHCHCH₂B⁻(C₆F₅)₃;

phenylmethylsilanediylbis(2-methyl-4,6-diisopropylindenyl)Zr+CH₂CHCHCH₂B-(C₆F₅)₃;

phenylmethylsilanediylbis(2-methyl-4-naphthylindenyl)Zr+ CH₂CHCHCH₂B⁻(C₆F₅)₃;

ethylenebis(2-methylindenyl) $Zr^+CH_2CHCHCH_2B^-(C_6F_5)_3$; ethylenebisindenyl $Zr^+CH_2CHCHCH_2B^-(C_6F_5)_3$;

ethylenebis(2-methyl-4,5-benzoindenyl)Zr⁺ CH₂CHCHCH₂B⁻(C₆F₅)₃;

ethylene(2-methyl-4,5-benzoindenyl)(2-methylindenyl)Zr⁺ CH₂CHCHCH₂B⁻(C₆F₅)₃;

ethylene(2-methyl-4,5-benzoindenyl)(2-methyl-4phenylindenyl)Zr+CH₂CHCHCH₂B-(C₆F₅)₃;

ethylene(2-methylindenyl)(4-phenylindenyl)Zr+ CH₂CHCHCH₂B⁻(C₆F₅)₃;

ethylenebis(2-methyl-4,5-benzoindenyl)Zr+ CH₂CHCHCH₂B-(C₆F₅)₃;

25 ethylenebis(2-methyl-4-phenylindenyl) Zr^+ $CH_2CHCHCH_2B^-(C_6F_5)_3$;

ethylenebis(2-methyl-4,6-diisopropylindenyl)Zr+ CH₂CHCHCH₂B-(C₆F₅)₃;

ethylenebis(2-methyl-4-naphthylindenyl) Zr^+ CH₂CHCHCH₂B⁻(C₆F₅)₃;

ethylenebis(2-ethyl-4-phenylindenyl) $Zr^+CH_2CHCHCH_2B^-$ (C_6F_5)₃;

ethylenebis(2-ethyl-4,6-diisopropylindenyl)Zr⁺ CH₂CHCHCH₂B⁻(C₆F₅)₃;

ethylenebis(2-ethyl-4-naphthylindenyl) Zr^+ CH₂CHCHCH₂B⁻(C₆F₅)₃;

dimethylsilanediylbis(2-ethyl-4-phenylindenyl)Zr⁺ CH₂CHCHCH₂B⁻(C₆F₅)₃;

dimethylsilanediylbis(2,3,5-trimethylcyclopentadienyl)Zr⁺ CH₂CHCHCH₂B⁻(C₆F₅)₃;

1,6-{bis[methylsilylbis(2-methyl-4-phenylindenyl)Zr+CH₂CHCHCH₂B-(C₆F₅)₃]}hexane;

45 1,6-{bis[methylsilylbis(2-ethyl-4-phenylindenyl)Zr+ CH₂CHCHCH₂B⁻(C₆F₅)₃]}hexane;

1,6-{bis[methylsilylbis(2-methyl-4-naphthylindenyl)Zr+CH₂CHCHCH₂B-(C₆F₅)₃]}hexane;

1,6-{bis[methylsilylbis(2-methyl-4,5-benzoindenyl)Zr+CH₂CHCHCH₂B-(C₆F₅)₃]}hexane;

1,6-{bis[methylsilyl(2-methyl-4-phenylindenyl)(2-methylindenyl)Zr+CH₂CHCHCH₂B-(C₆F₅)₃]}hexane;

1,2-{bis[methylsilylbis(2-methyl-4-phenylindenyl)Zr+CH2CHCH2B-(C₆F₅)₃]}ethane;

1,2-{bis[methylsilylbis(2-ethyl-4-phenylindenyl)Zr+ CH₂CHCHCH₂B-(C₆F₅)₃]}ethane;

1,2-{bis[methylsilylbis(2-methyl-4-naphthylindenyl)Zr+CH-CHCH_B-(C₆F₅)₃]}ethane;

 $CH_2CHCHCH_2B^-(C_6F_5)_3$] ethane; 1,2-{bis[methylsilylbis(2-methyl-4,5-benzoindenyl)Z⁺

CH₂CHCHCH₂B⁻(C₆F₅)₃]ethane;

1,2-{bis[methylsilyl(2-methyl-4-phenylindenyl)(2-methylindenyl)Zr+CH₂CHCHCH₂B-(C₆F₅)₃]}ethane.

The preparation of the novel transition metal compound of the formula I is illustrated by the following reaction schemes.

L, n, M, X, B, A, R¹ and m in the formulae II, III and IV are as defined in formula I and Hal is a halogen atom such as fluorine, chlorine, bromine or iodine.

Compounds of the formula II are described in the literature (J. Okuda, Topics in Current Chemistry, Vol. 160; Springer Verlag, Berlin Heidelberg 1991, page 97). Reaction of compounds of the formula II in inert solvents with dianion compounds of the formula III, for example 1,4butanediyllithium or 2-butene-1,4-diylmagnesium, leads to 20 in the polymerization. elimination of a salt and formation of the cyclic systems IV in which the M-X or the M-B bond can be a covalent bond or a coordination of the compound X-B to the metal

The compound of the formula IV can be reacted with 25 Lewis acids of the formula AR1 in organic solvents, for example toluene, benzene, methylene chloride, carbon tetrachloride and petroleum spirit, to give the compound of the

The novel transition metal compounds of the formula I 30 can be isolated or directly used for further reactions. The compounds of the formula I can also be prepared without isolation of intermediate and end stages in a single-vessel reaction from metallocene dihalides, dianion compounds

Suitable solvents for this purpose are aliphatic or aromatic solvents, such as hexane or toluene, or halogenated hydrocarbons, such as methylene chloride, or halogenated aromatic hydrocarbons such as o-dichlorobenzene.

A further possibility for preparing the novel compounds 40 of the formula I comprises the formation of metallocycles of the formula IV by electrocyclic ring-closure reaction of, for example, metallocene-bisolefin complexes or metalloceneolefin-aldehyde complexes and subsequent reaction with Ar^{1}_{m} .

The present invention also relates to a process for preparing an olefin polymer by polymerization of at least one olefin in the presence of a catalyst containing at least one transition metal compound of the formula I. The polymer-

Preference is given to polymerizing olefins of the formula R^a —CH=CH— R^b , where R^a and R^b are identical or different and are each a hydrogen atom, a halogen atom, an alkoxy group, a hydroxy group, an alkylhydroxy group, an aldehyde group, a carboxylic acid group or a carboxylic 55 ester group or a saturated or unsaturated hydrocarbon radical having from 1 to 20 carbon atoms, in particular from 1 to 10 carbon atoms, which can be substituted by an alkoxy group, a hydroxy group, an alkylhydroxy group, an aldehyde group, a carboxylic acid group or a carboxylic ester group, or Ra and R^b together with the atoms connecting them can form a one or more rings. Examples of such olefins are 1-olefins such as ethylene, propylene, 1-butene, 1-hexene, 4-methyl-1-pentene, 1-octene, styrene, cyclic olefins such as norbornene, vinylnorbornene, tetracyclododecene, 65 ethylidenenorbornene, dienes such as 1,3-butadiene or 1,4hexadiene, biscyclopentadiene or methyl methacrylate. In

particular, propylene or ethylene are homopolymerized, ethylene is copolymerized with one or more C₃-C₂₀-1-olefins, in particular propylene, and/or one or more C4-C20-dienes, in particular 1,3-butadiene, or norbornene and ethylene are copolymerized.

The polymerization is preferably carried out at a temperature of from -60 to 300° C., particularly preferably from 30 to 250° C. The pressure is from 0.5 to 2500 bar, preferably from 2 to 1500 bar. The polymerization can be carried out continuously or batchwise, in one or more stages, in solution, in suspension, in the gas phase or in a supercritical medium.

It is also possible to use mixtures of two or more transition metal compounds of the formula I. By this means, polyole-15 fins having a broad or multimodal molecular weight distribution can be obtained.

A prepolymerization can be carried out by means of the compound of the formula I. For the prepolymerization. preference is given to using the (or one of the) olefin(s) used

The compounds of the formula I can also be applied to a support, particularly for controlling the particle morphology. Suitable support materials are, for example, silica gels, aluminum oxides, solid aluminoxane or other inorganic support materials such as, for example, magnesium chloride. Another suitable support material is a polyolefin powder in finely divided form.

The supported catalyst system can be resuspended as powder or still together with solvent and can be metered into the polymerization system as a suspension in an inert suspension medium.

To remove catalyst poisons present in the olefin, purification using an aluminum alkyl, for example trimethylaluminum, triethylaluminum and Lewis acids and be directly used for the polymerization. 35 triisobutylaluminum, is advantageous. This purification can be carried out either in the polymerization system itself or the olefin is contacted with the Al compound and subsequently separated again prior to its addition to the polymerization system.

Hydrogen is added, if required, as molecular weight regulator and/or to increase the activity. The total pressure in the polymerization system is from 0.5 to 2500 bar, preferably from 2 to 1500 bar.

The compound of the formula I is here used in a 45 concentration, based on the transition metal, of preferably from 10^{-3} to 10^{-8} , particularly preferably from 10^{-4} to 10^{-7} mol of transition metal per dm3 of solvent or per dm3 of reactor volume.

If the polymerization is carried out as a suspension or ization can be a homopolymerization or a copolymerization. 50 solution polymerization, an inert solvent customary for the Ziegler low-pressure process is used. For example, the polymerization is carried out in an aliphatic or cycloaliphatic hydrocarbon; examples of such solvents which may be mentioned are propane, butane, hexane, heptane, isooctane, cyclohexane, methylcyclohexane. Furthermore, a petroleum or hydrogenated diesel oil fraction can be used. It is also possible to use toluene. Preference is given to carrying out the polymerization in the liquid monomer.

If inert solvents are used, the monomers are metered in in gaseous or liquid form.

The duration of the polymerization can be as desired. since the catalyst system to be used according to the invention has only a small fall in polymerization activity with

Prior to addition of the catalyst, in particular the supported catalyst system (containing at least one novel compound of the formula I, support material and/or a polyolefin powder in finely divided form), another aluminum alkyl compound such as, for example, trimethylaluminum, triethylaluminum, triisobutylaluminum, trioctylaluminum or isoprenylaluminum can additionally be added to the reactor to stabilize the polymerization system (for example for removing catalyst 5 poisons present in the olefin). This is added to the polymerization system in a concentration of from 100 to 0.01 mmol of Al per kg of reactor contents. Preference is given to triisobutylaluminum and triethylaluminum in a concentration of from 10 to 0.1 mmol of Al per kg of reactor contents. 10 This enables a small Al/M molar ratio to be selected in the synthesis of a supported catalyst system.

The novel compounds of the formula I are highly active catalyst components for olefin polymerization.

In principle, the use of cocatalysts in the polymerization 15 reaction is not required, i.e. the novel compounds of the formula I can be used as catalyst for olefin polymerization without the need for a cocatalyst such as aluminoxane.

The following examples serve to illustrate the invention.

General procedures: Preparation and handling of organometallic compounds were carried out with exclusion of air and moisture under argon protection (Schlenk technique).

All solvents required were dried prior to use by boiling for a number of hours over a suitable desiccant and subsequent distillation.

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The compounds were characterized using ¹H-NMR, ¹³C-NMR and IR spectroscopy.

A. Synthesis of the Compounds of the Formula IV

The preparation of the butadiene complexes was carried out according to G. Erker, K. Engel, Ch. Sarter in R. B. 30 King, J. J. Eisch, Organometallic Synthesis, Vol 3, Academic Press, New York 1986, 529:

EXAMPLE 1

Dimethylsilanediylbis(2-methyl-4,5-benzoindenyl) zirconium(η⁴-butadiene)

100 ml of toluene (precooled to -40° C.) are added to a mixture of 5.0 g (8.67 mmol) of dimethylsilanediylbis(2-methyl-4,5-benzoindenyl)zirconium dichloride and 2.3 g 40 (10.4 mmol, 1.2 equivalents) of (2-butene-1,4-diyl) magnesium bistetrahydrofuran ("butadienemagnesium") which has been cooled to -40° C. While stirring vigorously, the mixture is allowed slowly to warm up to room temperature. After stirring for four hours, the dark red solution is 45 separated via a frit from unreacted butadienemagnesium and magnesium chloride formed. The filtrate is evaporated to dryness and the residue is washed with 10 ml of pentane. This gives 4.4 g (70%) of a deep red powder.

EXAMPLE 2

$\begin{array}{c} \mbox{Dimethylsilanediylbis}(2\text{-methyl-4-phenylindenyl}) \\ \mbox{zirconium}(\eta^{4}\text{-butadiene}) \end{array}$

100 ml of toluene (precooled to -40° C.) are added to a 55 mixture of 5.0 g (7.95 mmol) of dimethylsilanediylbis(2-methyl-4-phenylindenyl)zirconium dichloride and 2.1 g (9.5 mmol, 1.2 equivalents) of (2-butene-1,4-diyl)magnesium bistetrahydrofuran ("butadienemagnesium") which has been cooled to -40° C. While stirring vigorously, the mixture is 60 allowed slowly to warm up to room temperature. After stirring for four hours, the dark red solution is separated via a frit from unreacted butadienemagnesium and magnesium chloride formed. The filtrate is evaporated to dryness and the residue is washed with 10 ml of pentane. This gives 3.5 g 65 (72%) of a red-brown powder.

B. Synthesis of the Compounds of the Formula I

10 EXAMPLE 3

Dimethylsilanediylbis(2-methyl-4,5-benzoindenyl) Zr⁺CH₂CHCHCH₂B⁻(C₆F₅)₃;

3.0 g (5.35 mmol) of dimethylsilanediylbis(2-methyl-4, 5-benzoindenyl)zirconium(η^4 -butadiene) are dissolved in 100 ml of toluene and admixed with 3.01 g (5.88 mmol, 1.1 equivalents) of tris(pentafluorophenyl)borane. The reaction solution is left stirring for 24 hours at room temperature and the strongly dark brown suspension is subsequently evaporated to half its volume. The precipitate is filtered off and washed with 10 ml of pentane. This gives 5.27 g (92%) of a sparingly soluble reddish brown powder.

EXAMPLE 4

Dimethylsilanediylbis(2-methyl-4-phenylindenyl) $Zr^+CH_2CHCHCH_2B^-(C_6F_5)_3$;

3.0 g (4.90 mmol) of dimethylsilanediylbis(2-methyl-4-phenylindenyl)zirconium(η^4 -butadiene) are dissolved in 100 ml of toluene and admixed with 2.76 g (5.39 mmol, 1.1 equivalents) of tris(pentafluorophenyl)borane. The reaction solution is left stirring for 24 hours at room temperature and the strongly dark brown suspension is subsequently evaporated to half its volume. The precipitate is filtered off and washed with 10 ml of pentane. This gives 4.84 g (88%) of a sparingly soluble reddish brown powder.

C. Polymerization Examples

EXAMPLE 5

The catalyst solution is prepared by mixing 11 mg of biscyclopentadienylzirconium(η^4 -butadiene) in 20 ml of toluene with a solution of 20.4 mg of tris(pentafluorophenyl) borane in 20 ml of toluene. 900 ml of toluene are admixed with 1 ml of a 10% strength by weight TIBA solution in toluene and subsequently with 1 ml of this catalyst solution.

For the polymerization, this solution is placed in an inert $1.5 \, \mathrm{dm^3}$ stirred reactor, heated to 70° C. and polymerization is carried out at a pressure of 7 bar of ethylene. After 2 hours, the reactor is vented, the polymer is filtered from the suspension, washed with acetone and dried for 12 hours in a vacuum drying oven. This gives 38 g of polyethylene having an M_{w} of 297,000 g/mol and an M_{w}/M_{n} of 2.5 according to GPC.

EXAMPLE 6

The polymerization of Example 5 is repeated using highpurity ethylene, except that no TIBA was added. This gives 37 g of polyethylene having identical properties.

EXAMPLE 7

The polymerization of Example 5 is repeated, except that 2 ml of the catalyst solution are added and 100 ml of 1-hexene were first added to the reactor and 5 bar of ethylene were subsequently added. After 30 minutes, the reactor is vented, the polymerization is stopped using methanol and the polymer is filtered from the suspension, washed with acetone and dried for 12 hours in a vacuum drying oven. This gives 25 g of an ethylene/1-hexene copolymer containing 5.2 mol % of hexene (according to ¹³C-NMR) and having an M_w of 60,000 g/mol and an M_w/M_n of 2.6 according to GPC. The DSC melting point of the 2nd heating is 110° C.

EXAMPLE 8

A 1.5 dm³ autoclave, which was thoroughly flushed with ethene beforehand, is charged with 600 cm³ of an 85%

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strength by weight solution of norbornene in toluene. The solution is saturated with ethene by repeated pressurization with ethene (18 bar). A suspension of 2.28 mg of 4- $(\eta^5$ cyclopentadienyl)-4,7,7-trimethyl-(η⁵-4,5,6,7tetrahydroindenyl)Zr+CH₂CHCHCH₂B-(C₆F₅)₃ in 10 ml of 5 toluene was added to the reactor thus prepared (in the case of hydrogen regulation, hydrogen can be injected at this point). Polymerization was carried out for one hour while stirring, with the ethene pressure being kept at 18 bar by metering in further amounts.

After the end of the reaction time, the polymerization mixture was drained into a vessel and immediately introduced into 5 dm3 of acetone, the mixture was stirred for 10 min and the precipitated product was subsequently filtered off. The filter cake was washed alternately three times each 15 with 10% strength hydrochloric acid and acetone. It was subsequently washed to neutrality with water, the residue was slurried in acetone and filtered again. The polymer thus purified was dried at 80° C. for 15 hours in vacuo (0.2 bar). After drying, there were obtained 224 g of colorless polymer 20 which had a glass transition temperature of 179° C., a viscosity number of 52 cm³/g, a tensile strength of 59 MPa and an elongation at break of 3.1%. The activity was 80.5 kg of polymer/h×mmol.

EXAMPLE 9

106 mg (0.385 mmol) of biscyclopentadienylzirconium (η⁴-butadiene) were dissolved in toluene and admixed with a solution of 186 mg (0.363 mmol) of $B(C_6F_5)_3$ in toluene. The formation of the catalyst can be recognized by the appearance of turbidity or of a precipitate. In parallel thereto, a dry 16 dm3 reactor was flushed first with nitrogen and subsequently with propylene and then charged with 10 dm³ of liquid propylene. 15 cm³ of triisobutylaluminum (20% strength in hydrocarbon, 12 mmol) were then added to the reactor and the mixture was stirred at 30° C. for 15 minutes. The catalyst suspension was subsequently added to the reactor, heated to the polymerization temperature of 70° C.(4° C./min) and the polymerization system was held for 1 hour at 70° C. by means of cooling. The polymerization was stopped by addition of 20 ml of isopropanol. The excess monomer was vented and residues of solvent were taken off in vacuo. This gave 850 g of liquid, atactic polypropylene. The catalyst activity was 8 kg of PP/(g of metallocenexh). 45

 $VN=5 \text{ cm}^3/\text{g}, M_w=1500 \text{ g/mol}, M_w/M_n=3.2.$

EXAMPLE 10

10 mg (17.9 μmol) of rac-dimethylsilanediylbis(2methyl-4,5-benzo-1-indenyl)zirconium(η⁴-butadiene) were ⁵⁰ dissolved in 10 ml of toluene and admixed with 10 ml of a solution of 9.2 mg (18 mmol) of B(C₆F₅)₃ in toluene. The formation of the catalyst can be recognized by the appearance of turbidity or of a dark precipitate. The polymerization was carried out by a method similar to Example 9 at 70° C. 55 The excess monomer was vented and the polymer powder was dried in vacuo. This gave 2500 g of isotactic polypro-

The catalyst activity was 250 kg of PP/(g of metallocenex

VN=240 cm³/g, mp.=148.7° C., M_w=298,000 g/mol, $M_w/M_n=2.2$, $MFI_{(230/2.16)}=3.2$ dg/min.

EXAMPLE 11

The preparation of the catalyst suspension from Example 10 was repeated, except that 3 mg (4.8 μ mol) of rac-

dimethylsilanediylbis(2-methyl-4-phenyl-1-indenyl) zirconium(η⁴-butadiene) dissolved in 10 cm³ of toluene were reacted with 2.5 mg (4.9 μ mol) of B(C₆F₅)₃ dissolved in 10 cm³ of toluene. The polymerization was carried out at 60° C. after adding the catalyst suspension to the reactor. The polymerization gave 2250 g of isotactic polypropylene

The catalyst activity was 750 kg of PP/(g of metallocenex h).

VN=620 cm³/g, mp.=155° C., MFI_(230/5)=0.35 dg/min.

EXAMPLE 12

The preparation of the catalyst suspension from Example 10 was repeated, except that 10 mg (21 µmol) of racdimethylsilanediylbis(2-methyl-1-indenyl)zirconium(n⁴butadiene) dissolved in 10 cm³ of toluene were reacted with 10.7 mg (21 μ mol) of B(C₆F₅)₃ dissolved in 10 cm³ of toluene. The polymerization gave 1900 g of isotactic polypropylene powder. The catalyst activity was 190 kg of PP/(g of metallocenexh).

VN=180 cm³/g, mp.=145° C., M_w =192,000, M_w/M_n =2.2, MFI_(230/2.16)=12 dg/min.

EXAMPLE 13

The preparation of the catalyst suspension from Example 10 was repeated, except that 10 mg (20.2 µmol) of phenylmethylmethylenefluorenylcyclopentadienylzirconium-(\(\eta^4\)-butadiene) dissolved is 10 cm³ of toluene were reacted with 10.7 mg (21 μ mol) of B(C₆F₅)₃ dissolved in 10 cm³ of toluene. The polymerization gave 1100 g of syndiotactic polypropylene powder.

The catalyst activity was 110 kg of PP/(g of metallocenex h).

 $VN=137 \text{ cm}^3/g$, mp.=133° C., $M_w=122,000 \text{ g/mol}$, $M_{w}/M_{n}=2.3$.

EXAMPLE 14

Reactor Preparation

A 1500 ml reactor which had been repeatedly flushed with nitrogen was charged with 1000 ml of an 85% strength by weight solution of norbornene in toluene and the solution was heated to 70° C. The solution was saturated with ethylene by repeated pressurization with ethylene (16 bar gauge pressure). 2 ml of a 20% strength triethylaluminum solution in toluene were added to the depressurized reactor in a countercurrent of ethylene and the mixture was subsequently stirred for 15 minutes.

Catalyst Preparation

A solution of 12.6 mg of tris(pentafluorophenyl)borane (0.025 mmol) in 1.26 ml of toluene was added to the solution of 9.0 mg of (butadiene)isopropylene(1-indenyl) cyclopentadienylzirconium (0.025 mmol) in 0.9 ml of toluene. The catalyst mixture was preactivated for 25 minutes at room temperature.

Polymerization and Isolation

The catalyst mixture was added to the prepared reaction solution and an ethylene pressure of 16 bar was subsequently applied as quickly as possible. Polymerization was carried out for 2 hours at 70° C. while stirring at about 750 rpm, with the ethylene pressure being kept constant at 16 bar.

To end the reaction, the reactor was first vented and the reaction solution was subsequently drained into a vessel. The polymer was precipitated in 2500 ml of acetone and filtered after stirring for 5 minutes. The filter cake was repeatedly washed alternately with 10% strength hydrochloric acid and acetone. It was subsequently washed to neutrality with water and, after addition of 1000 ml of acetone, filtered. The powder thus purified was dried for 15 hours at 80° C. at a pressure of 0.2 bar.

Characterization

After drying, 30 g of colorless powder were obtained; this corresponds to an activity of 610 g of polymer/hxmmol of metallocene. A viscosity number of 106 cm³/g and a glass transition temperature of 135° C. were measured on the polymer. No melting point was able to be detected by means 10 of thermal analysis.

EXAMPLE 15

30 mg (31 µmol) of rac-dimethylsilanediylbis(2-methyl- 15 1-indenyl)Zr+CH2CHCHCH2B-(C6F5)3 are dissolved in 30 ml of toluene. 10 g of SiO₂ are stirred into the catalyst solution until a homogeneous distribution is achieved.

The solid is then filtered off from the solvent and washed twice with 10 ml of toluene.

In parallel thereto, a dry 16 dm3 reactor was flushed first with nitrogen and subsequently with propylene and charged with 10 dm3 of liquid propylene. 15 cm3 of triisobutylaluminum (20% strength in hydrocarbon, 12 mmol) were then added to the reactor and the mixture was stirred at 30° C. for 15 minutes. The catalyst suspension was subsequently added to the reactor, heated to the polymerization temperature of 70° C. (4° C./min) and the polymerization system was held stopped by addition of 20 ml of isopropanol. The excess monomer was vented and solvent residues were taken off in

The polymerization gives 2.5 kg of isotactic polypropylene powder. The catalyst activity was 83 kg of PP/(g of 35 M_w/M_n=2.1 metallocenexh).

 $VN=184 \text{ cm}^3/\text{g}$, mp.=145° C., $M_{w}=193,000 \text{ g/mol}$, $M_{u}/M_{u}=2.2.$

The reactor shows no deposits on walls or stirrer.

EXAMPLE 16

a) Preparation of the Catalyst Component

A solution of 1 µmol of biscyclopentadienylzirconium 45 (n⁴-butadiene) in 1 ml of toluene is admixed with a solution of 1 µmol of tris(pentafluorophenyl)borane in 10 ml of toluene and, after a reaction time of 15 minutes, evaporated in vacuo to a volume of 2 ml. In parallel thereto, 4 g of RAccurel LDPE powder in a sieve fraction of smaller than 50 200 um are dried in vacuo and flushed with argon. The support powder is stirred into the catalyst solution until a homogeneous distribution is achieved.

b) Polymerization

A dry 1.5 dm³ stirred reactor is flushed with nitrogen to remove the oxygen and charged with 0.9 dm3 of an inert diesel oil (bp. from 100 to 120° C.). After flushing with ethylene, it is heated to 70° C. and the catalyst is added as powder. Polymerization is subsequently carried out without 60 additional activator at a pressure of 7 bar of ethylene. After 1 hour, the reactor is vented, the polymer is filtered from the suspension and dried for 12 hours in a vacuum drying oven. This gives 18 g of polyethylene powder having a bulk density of 0.253 kg/dm³ and a viscosity number VN of 389 cm³/g. The polydispersity M_u/M_u is 2.6 (according to GPC). The reactor shows no deposits on walls or stirrer.

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EXAMPLE 17

a) Preparation of the Catalyst Component

BiscyclopentadicnylZr+CH2CIICIICH2B-(C6F5)3 was 5 supported on SiO₂ using a method similar to Example 15.

b) Gas-Phase Polymerization of Ethylene

A gas-phase polymerization of ethylene was carried out in a 2 dm³ steel autoclave having polished walls. The fluidized bed was generated mechanically by means of a double-helix stirrer going around the wall, using an initial charge of 10 g of polyethylene powder as seed bed. Via a pressure burette, first the cocatalyst (2 mmol) of triisobutylaluminum in 2 cm³ of isopentane and subsequently 1 g of the catalyst mixture (19.2 umol of Zr) were metered in. The polymerization was subsequently carried out at 8 bar partial pressure of ethylene at a temperature of 80° C. for 1 hour and ended by venting the autoclave.

This gave 118 g of polyethylene having a VN of 309 ml/g, corresponding to an activity of 6.2 kg of PE/mmol of metallocene.

EXAMPLE 18

The preparation of the catalyst suspension of Example 10 25 was repeated, except that 10 mg (24 μ mol) of dimethylmethylene-9-fluorenylcyclopentadienylzirconium (4-butadiene) dissolved in 10 cm³ of toluene were reacted with 12.8 mg (25 μ mol) of B(C₆F₅)₃ dissolved in 10 cm³ of toluene. The polymerization was carried out after metering at 70° C. for 1 hour by cooling. The polymerization was 30 the catalyst suspension into the reactor at 60° C. The polymerization gave 900 g of syndiotactic polypropylene powder. The catalyst activity was 90 kg of PP/(g of metallocenexh).

 $VN=92 \text{ cm}^3/\text{g}, \text{ m.p.}=126^{\circ} \text{ C., } M_{\text{w.}}=63,000 \text{ g/mol,}$

EXAMPLE 19

The preparation of the catalyst suspension of Example 10 was repeated, except that 5 mg (8 μ mol) of racdimethylsilanediylbis(2-methyl-4,6-diisopropyl-1-indenyl) zirconium(4-butadiene) dissolved in 10 cm³ of toluene were reacted with 4.1 mg (8.1 µmol) of B(C₆F₅)₃ dissolved in 10 cm3 of toluene. The polymerization was carried out after metering the catalyst suspension into the reactor at 60° C. The polymerization gave 2100 g of isotactic polypropylene powder.

The catalyst activity was 420 kg of PP/(g of metallocenex

 $VN=423 \text{ cm}^3/\text{g}, \text{ mp.=}154^\circ \text{ C., MFI}_{(230/5)}=3.1 \text{ dg/min,}$ $M_w=588,000 \text{ g/mol}, M_w/M_n=3.5.$

EXAMPLE 20

The preparation of the catalyst suspension of Example 11 was repeated. The polymerization was carried out after metering the catalyst suspension into the reactor at 70° C. The polymerization gave 2800 g of isotactic polypropylene powder.

The catalyst activity was 933 kg of PP/(g of metallocenex

VN=544 cm³/g, mp.=154° C., MFI_(230/5)=1.3 dg/min, $M_w = 741,000 \text{ g/mol}, M_w/M_n = 2.8.$

EXAMPLE 21

The preparation of the catalyst suspension of Example 12 was repeated, except that the 16 dm³ reactor was charged